THE RONDELETIA COMPLEX
(RUBIACEAE): AN ATTEMPT TO
USE ITS, RPS16, AND TRNL-F
SEQUENCE DATA TO DELIMIT
GUETTARDEAE, RONDELETIEAE,
AND SECTIONS WITHIN
RONDELETIA¹

Johan H. E. Rova,² Piero G. Delprete,³ and Birgitta Bremer⁴

ABSTŘACT

In the present study, a molecular phylogeny of the Rondeletia L. complex (Rubiaceae, Rondeletieae) was constructed with the following main objectives: (1) to evaluate the sections of Rondeletia proposed by Fernández Zequeira; (2) to test if Stevensia Poit. belongs to the Rondeletieae s. str.; (3) to check if ITS data from Rondeletieae support previous phylogenetic results from trnL-F data regarding circumscription of Rondeletieae; and (4) to verify if Hodgkinsonia F. Muell. belongs to Guettardeae or elsewhere. Two analyses were performed, one with ITS sequences from 46 taxa in the Rondeletieae–Guettardeae complex, and the other with combined ITS, rps16, and trnL-F sequences of 21 taxa. Representatives of nine of the 10 sections of Rondeletia recognized by Fernández Zequeira were included in the ITS analysis. Five of her sections could be tested for monophyly. Support was only found for Rondeletia sect. Leoninae M. Fernández Zeq., while representatives from section Chamaebuxifoliae M. Fernández Zeq., section Hypoleucae Standl., and section Nipenses M. Fernández Zeq. together form a well-supported clade that could be distinguished also based on morphology. The latter clade is sister to Stevensia, which is thus placed within Rondeletia s. str. In addition, ITS sequence data confirm the separation of Rovaeanthus Borhidi from Rogiera Planch. Support is low for inclusion of Blepharidium Standl., Mazaea Krug & Urb., Phyllomelia Griseb., Rachicallis DC., part of Rogiera, and Suberanthus Borhidi & M. Fernández Zeq. in Rondeletieae. Rachicallis, Mazaea, and Phyllomelia form a clade with strong support. The tribe Rondeletieae s. str. was found to be monophyletic in all trees, although with low support; however, a redelimitation of the tribe is proposed here based on this study and previous phylogenetic analyses. The monophyly for the tribe Guettardeae is weakly supported, with the inclusion of Arachnothryx Planch. (including Cuatrecasasiodendron Steverm.), Gonzalagunia Ruiz & Pav., Hodgkinsonia, and Timonius DC. Although it was recently the subject of a molecular phylogenetic study, the tribe Guettardeae is still in need of a wide-ranging survey in order to confirm its monophyly and delimit its taxonomic boundaries. Because Cuatrecasasiodendron was found within the Arachnothryx clade, the two genera are here synonymized as Arachnothryx, and in turn positioned within the tribe Guettardeae. In addition, based on herbarium and field studies, the two species described under Cuatrecasasiodendron (C. spectabile Steyerm, and C. colombianum Standl. & Steyerm.) are treated as synonyms to the new combination Arachnothryx spectabilis (Steyerm.) Rova, Delprete & B. Bremer, which is proposed here.

Key words: Arachnothryx, Cuatrecasasiodendron, Guettardeae, Hodgkinsonia, ITS, phylogeny, Rogiera, Rondeletia, Rondeletieae, rps16, Rubiaceae, Stevensia, trnL-F.

doi: 10.3417/2006179

¹We are grateful to Attila Borhidi (Janos Pannonius University, Pécs) for help with identification of specimens of *Rondeletia* and related taxa and to Nahid Heidari and the staff at the Evolutionary Biology Centre (Uppsala University) for technical assistance. The National Tropical Botanical Garden, Lawai, Kauai, Hawaii, provided material for *Rondeletia inermis* and *Rovaeanthus strigosus*. This work was supported by a grant from the Bergius Foundation. Part of this research was realized during a Visiting Scientist fellowship to Piero Delprete (PD) from the Conselho Nacional de Desenvolvimento Científico e Tecnológico (CNPq) at the Universidade Federal de Goiás, Goiânia, Goiás, Brazil. PD's fieldwork in Cuba was supported by a grant from the MacArthur Foundation through the American Council of Learned Societies/Social Science Research Council Working Group on Cuba. We also thank the Cuban government for giving permission to PD to perform fieldwork, and Ramona Oviedo, Mayra Fernández, Jaqueline Salazar, and Lazara Sotolongo (Academia de Ciencias, Instituto de Ecología y Sistemática, La Habana, Cuba; HAC) for accompanying PD during fieldwork in 2002.

² Hagaberg Baskarp, SE-566 92 Habo, Sweden. Author for correspondence: jorova@telia.com.

³ CNPq Visiting Scientist, Institute of Biological Sciences—ICB-1, Department of General Biology/Botany, Universidade Federal de Goiás, Campus II, 74001-970 Goiânia, Goiás, Brazil. Current address: Institut de Recherche pour le Développement, Botanique et Bioinformatique de l'Architecture des Plantes (AMAP), TA-A51/PS2, Blvd. de la Lironde, 34398 Montpellier Cedex 5, France. pdelprete@hotmail.com.

⁴ Bergius Foundation at the Royal Swedish Academy of Sciences and Botany Department, Stockholm University, SE-106 91 Stockholm, Sweden.

The tribe Rondeletieae (Rubiaceae, Cinchonoideae) includes predominantly shrubs and trees and is mostly distributed in the New World tropics (Robbrecht, 1988; Delprete, 1999a), with the main center of diversity in the Greater Antilles. A thorough description of the taxonomic and systematic history of the tribe is found in Delprete (1999a).

The largest genus of the tribe, Rondeletia L., is mainly Antillean and comprises approximately 120 species. Standley (1918) divided Rondeletia into 15 sections based on morphological and distributional data. Since then, several morphological and molecular studies in the Rondeletieae have argued about the status of Standley's sections and the circumscription of the genus *Rondeletia*. One opinion is that Rondeletia should be regarded as a narrowly circumscribed genus, separated from morphologically similar genera such as Arachnothryx Planch., Javorkaea Borhidi & Jarai-Koml., Rogiera Planch., Roigella Borhidi & M. Fernández Zeq., Rovaeanthus Borhidi, and Suberanthus Borhidi & M. Fernández Zeq. (Steyermark, 1967; Borhidi & Fernández Zequeira, 1981a, b; Borhidi, 1982, 1989, 1994; Borhidi & Járai-Komlódi, 1983; Fernández Zequeira, 1994; Delprete, 1999a, as Rondeletia complex sensu Delprete; Rova, 1999; Rova et al., 2002; Borhidi et al., 2004; Rova, unpublished). On the other hand, Lorence (1991) recognized Rondeletia as a widely circumscribed genus, treating the names applied to Mexican and Central American taxa of the complex as synonyms. Based on morphological data, Fernández Zequeira (1994) made an attempt to classify the Greater Antillean (especially the Cuban) Rondeletia species into 10 sections. Her classification comprised 104 species, most of them endemic to Cuba. This means that a majority of the species of *Rondeletia* s. str. were included in her study. According to Fernández Zequeira (1994), the sections are distinguished by various combinations of (often multistate) morphological characters such as position and shape of inflorescence, flower merosity, calyx lobe shape, leaf indumentum, and phyllotaxy (leaves opposite vs. verticillate). However, her focus on Cuban species did little to resolve the problem in the larger Rondeletia complex. The first aim of the present study was to test if Fernández Zequeira's sections of Rondeletia are supported by phylogenies obtained from molecular sequences. The second aim was to test if Stevensia Poit. was closely related to Rondeletia or not. Stevensia has not been included previously in molecular phylogenies, but morphology suggests a close affinity between the genera. Earlier studies (Bremer et al., 1995; Bremer & Thulin, 1998; Andersson & Rova, 1999; Rova et al., 2002) have shown that the tribes Guettardeae and Rondeletieae

are closely related, and this study also aimed to investigate if ITS data would suggest a similar circumscription of Rondeletieae as previous studies had. Finally, the study was aimed to investigate if ITS sequence data would support Bremer's (1992) inclusion of the Australian genus *Hodgkinsonia* F. Muell. in the Guettardeae or Delprete's (1996) transfer of the genus from the Guettardeae to the Chiococceae.

MATERIALS AND METHODS

For the ITS analyses, material was sampled from as many Rondeletia species and subspecies as possible. An effort was made to include representatives from all genera in Rondeletieae sensu Rova et al. (2002). The outgroup consisted of Luculia Sweet (basal in Rubiaceae, e.g., Bremer et al., 1995), Catesbaea fuertesii Urb., Chiococca alba (L.) Hitche. (Chiococcae s.l.), and 12 accessions representing 11 species in the following six genera of the tribe Guettardeae (based on available material and the results from Rova et al., 2002): Arachnothryx, Cuatrecasasiodendron Steyerm., Gonzalagunia Ruiz & Pav., Guettarda L., Rogiera, and Timonius DC. Authors of species names are given in Table 1, or otherwise when first mentioned in the text.

Fresh or silica gel—dried leaves were used for DNA extraction when available, but often herbarium material had to be used. DNA was extracted using the CTAB method (Doyle & Doyle, 1987) and cleaned with the QIAquick PCR Purification Kit (QIAGEN GmbH, Hilden, Germany). The cocktail for polymerase chain reaction (PCR) amplification was mixed as follows (to ca. 25 μ l): 2.5 μ l 10 \times buffer, 2.5 μ l 25 mM MgCl₂, 2 μl dNTP, 0.125 μl Taq DNA polymerase, 0.625 μl 10 μM forward primer, 0.625 μl 10 μM reverse primer, 2.5 μl 0.1 M TMACl, 2 μl dimethyl sulfoxide (DMSO), 2 µl template, and 10 µl water. In some cases, the amount of primer or template was doubled (replacing some of the water). Primers P17 and 26S-82R (Popp & Oxelman, 2001) were used for amplification. Sequencing reactions were realized using the DYEnamic ET terminator Cycle Sequencing kit (Amersham Biosciences, Buckinghamshire, England) following the protocol of the manufacturer (DMSO was added in the same concentration as in the PCR mix) and run on a MegaBACE 1000 DNA Analysis System (Amersham Biosciences). For sequencing, the same primers were used as in the PCR amplification.

For the ITS study, 50 new ITS sequences were produced, and five additional sequences were downloaded from GenBank and included in the data matrix. Taxon names, authors, vouchers, and GenBank accession numbers are presented in Table 1.

Manual alignment and gapcoding of the ITS sequences were performed with the following criteria: (1) an effort was made to see if gaps/insertions could be interpreted as caused by repeats or inversions, and if so, sequences were aligned to fit these possible events; (2) gaps (i.e., inferred insertion/deletion events) were introduced into the sequences to keep the number of substitutions in an aligned region to a minimum; (3) insertions/deletions and substitutions were considered equally probable events; and (4) gaps/insertions of equal length shared by two or more taxa were inferred to be homologous and binary coded. Gaps of more than one position in length introduced due to multiplication of single nucleotides, e.g., poly-A, were not coded. Regions where alignment could not be unambiguously interpreted were removed from the analysis. After alignment, two ITS matrices were produced, one including gap codings and the other without them. Two parsimony analyses, conducted with PAUP* version 4.0b10 (Swofford, 2000), were performed for each matrix. The first ITS analysis was a heuristic search (random addition sequence with 1000 replicates, tree bisection-reconnection [TBR] branch swapping, and MULTREES option in effect), and the second analysis was a jackknife search (faststep search option, 10,000 replicates, and Jac resample emulation).

For the combined analyses, the data matrix from the ITS study was combined with the entire matrices from the trnL-F study of Rova et al. (2002) and from an rps16 analysis (Rova, unpublished), keeping the indel codings from each matrix. Previous analyses of each separate data set resulted in similar trees, which implied that the data sets were congruent. Taxa not included in the combined analyses were then deleted using the command DELETE in the PAUP block. The resulting set of sequences comprised 20 ingroup taxa. This set included all taxa where all three sequences were available and all Rondeletia species where at least ITS and rps16 sequences were available. Chiococca alba was used as outgroup, because, previously, it had been clearly shown not to be part of the ingroup (Rova et al., 2002). The data were analyzed by a heuristic search (random addition sequence with 1000 replicates, TBR branch swapping, and MULTREES option in effect).

RESULTS

More than 50 DNA extractions were obtained from Rondeletia representatives, but only 27 of these (representing 23 species) were amplified by PCR and yielded sequences. Extractions that did not produce usable sequences were mostly made from herbarium material more than 50 years old. Material collected in silica gel almost always worked for PCR

and resulted in high-quality sequences. It was not possible to obtain sequences from section *Lindenianae* M. Fernández Zeq., although extractions were attempted from two different specimens. It was also not possible to obtain PCR products from Roigella correifolia (Griseb.) Borhidi & M. Fernández Zeq., which trnL-F data showed to be closely related to Rondeletia s. str. (Rova et al., 2002). For four sections (Rondeletia sect. Odoratae Standl., section Pedicellares Standl., section Rigidae M. Fernández Zeq., and section Chamaebuxifoliae), it was only possible to sequence one species from each section. We were unfortunately not able to sequence the type species of Rondeletia, R. americana L. This species seems to be very rarely collected, and extractions made from the herbarium material that we found in the Swedish Museum of Natural History Herbarium did not amplify despite several attempts. We were not able to establish contact with anyone who could assist us with recently collected material from St. Vincent, where the species is endemic, and it was not possible to do such fieldwork ourselves. Ten species that yielded sequences were not listed under any section in the work of Fernández Zequeira, but four of them could be assigned to sections based on the key provided in her paper (Fernández Zequeira, 1994): R. inermis (Spreng.) Krug & Urb. and R. pilosa Sw. belonging to section Leoninae, and R. hameliifolia Dwyer & M. V. Hayden and R. purdiei Hook. f. belonging to section Calophyllae M. Fernández Zeq. Sectional affinities are indicated in Figure 1.

The first ITS matrix, without indel coding, included 699 characters, of which 174 were parsimony informative. The second ITS matrix, where indels were coded, included 723 characters, of which 198 were parsimony informative. The combined ITS, rps16, and trnL-F matrix included 2751 characters, 1451 of which were parsimony informative.

The strict consensus tree obtained from the ITS analyses is presented in Figure 1. Heuristic searches of both data sets each resulted in 48 most parsimonious trees. Tree lengths were 768 (consistency index [CI] = 0.56, retention index [RI] = 0.76) in the heuristic search of the data set without indel coding and 805 (CI = 0.65, RI = 0.77) in the heuristic search where indels were coded. Strict consensus trees were identical for both data sets. Jackknife support was not found for all clades in the strict consensus trees from the heuristic searches, and jackknife support values for a clade could vary up to more than 10 units between the two data sets. Tree topologies differed only marginally between the two jackknife searches. The jackknife search without indel codings found one clade that was not found in the other jackknife search (or in the heuristic

searches), and the jackknife search with indel codings resulted in two clades not found in the jackknife search without indels coded (Fig. 1). Support for these clades was low in all cases.

The analysis of the combined matrix resulted in 12 equally parsimonious trees (length 2046, CI = 0.91, RI = 0.97). A strict consensus of these trees is shown in Figure 2, and branches from the consensus tree that also occur in the ITS analysis are marked in bold in Figure 1.

DISCUSSION

SECTIONAL CLASSIFICATION OF RONDELETIA

The main aim of our study was to test Fernández Zequeira's (1994) classification with 10 sections of Rondeletia using molecular phylogenetic analyses. This goal was hard to reach satisfactorily; despite an extensive search, it was difficult to find herbarium or silica gel—dried material that would work for PCR and sequencing. For five of the sections, only one representative of each could be sequenced. Furthermore, ITS data are obviously not variable enough to provide resolution among sections Hypoleucae and Nipenses. Nevertheless, we obtained several interesting results with regard to the circumscription of Rondeletia and some of Fernández Zequeira's sections.

There is strong support for the *Rondeletia* s. str. clade (Fig. 1, clade E). This clade consists of predominantly Antillean species. The only exceptions to this distribution are *R. hameliifolia* from Central America (Panama) and *R. purdiei* from South America (Ecuador). Neither *R. hameliifolia* nor *R. purdiei* were included in Fernández Zequeira's (1994) treatment, but according to her identification key, both species would belong to section *Calophyllae*. In our study, the two species form a clade with strong support. A third representative of this section is *R. alaternoides* A. Rich. from Cuba, which is found in clade F (Fig. 1). Thus, ITS sequence data do not support a monophyletic section *Calophyllae*.

Rondeletia deamii (Donn. Sm.) Standl. is found just outside the Rondeletia s. str. clade. The generic position of this Central American species has recently been under debate. Lorence (1999) supported its position in Rondeletia, but Borhidi (2001a) positioned it in Arachnothryx. Our ITS sequence data suggest that this species should be treated as a Rondeletia, although support for this hypothesis is less than 50.

Rondeletia inermis and R. pilosa Sw. were not included in Fernández Zequeira's (1994) treatment of Cuban Rondeletia, as these species occur in Puerto Rico and the U.S. Virgin Islands, respectively. However, according to her key to sections, they would

both belong to *Rondeletia* sect. *Leoninae*. In our analysis, they form a clade with strong support. *Rondeletia* sect. *Leoninae* would thus be the only one of Fernández Zequeira's sections that is supported by our ITS sequence data.

In all analyses, there is moderate support for a clade with *Rondeletia alaternoides*, *R. odorata* Jacq., and *R. pachyphylla* Krug & Urb. (Fig. 1, clade F), which represent sections *Calophyllae*, *Odoratae*, and *Pedicellares*, respectively. Following the diagnostic table of sections in Fernández Zequeira (1994), we were unable to find any morphological characters that support this group.

Rondeletia intermixta Britton and R. ochracea Urb. form a clade with strong support. While R. intermixta belongs to section Rondeletia M. Fernández Zeq., R. ochracea has not been previously classified to any section. It is thus possible to argue that R. ochracea should also belong to section Rondeletia. The only other known representative of section Rondeletia included in our analysis, R. portoricensis Krug & Urb., is placed in an unresolved relationship to the R. intermixta—R. ochracea clade, although jackknife support for this is below 50.

Our study does not show any support for a separation of sections *Hypoleucae* and *Nipenses* in *Rondeletia*. All sequenced representatives of these sections, except *R. berteroana* DC., are found in a strongly supported but unresolved clade (Fig. 1, clade H). No morphological character combination seems to be unique for these two sections as one group, according to the character list in Fernández Zequeira (1994). *Rondeletia berteroana* differs from the other sequenced species of section *Hypoleucae* (and section *Nipenses*) in being from Hispaniola. This species is found as sister to clade H but with very low support (Fig. 1).

Rondeletia chamaebuxifolia Griseb., the only sequenced representative of section Chamaebuxifoliae, is found closely related to the species from sections Hypoleucae and Nipenses. Following the diagnostic characters provided in Fernández Zequeira (1994) for sections Chamaebuxifoliae, Hypoleucae, and Nipenses, this clade (Fig. 1, clade G) could be distinguished from other sections by having 1- to 3-flowered inflorescences and retrorse-pilose flowers.

STEVENSIA

The second aim of our study was to investigate the relationships between *Stevensia* and *Rondeletia*. *Stevensia* is here for the first time included in a molecular phylogenetic study. According to ITS data, there is strong support for an inclusion of at least *S. minutifolia* Alain in *Rondeletia* s. str. The genus

GenBank accession numbers for all publication. previous for both ITS and rps16 sequencing unless the other sequence is cited from a s well as Table of taxa included in the study, including ITS and rosl6 voucher information for sequences originally presented in this paper, a sequences used. If only one voucher is mentioned, the same voucher is used Table

Taxon	Voucher data	ITS	9Isd ₁	trnL-F
Acrosynanthus revolutus Urb.	Cuba, P. G. Delprete et al. 8818 (UPS)	AY730288		
4rachnothryx buddleioides (Benth.) Planch.	Panama, J. Rova & Sundbaum 2411 (S)	AY730299		
Arachnothryx chimboracensis (Standl.) Steyerm.	Ecuador, P. G. Delprete & A. Verduga 6398 (NY)	AY730292		
Arachnothryx leucophylla (Kunth) Planch.	Cuba, J. Rova et al. 2287 (GB)	AY730296	AF242910	AF1527186
Arachnothryx sp. indet.	Ecuador, G. Harling 27108 (NY)	AY730298		
Blepharidium guatemalense Standl.	Guatemala, Gustafsson et al. 212 (GB)	AY730287	AF242916	AF1527356
Catesbaea fuertesii Urb.		$\rm AY205364^{1}$		
Chiococca alba (L.) Hitche.		$\rm AY205367^{1}$	$AF004034^{4}$	$AF102400^{7}$
Cuatrecasasiodendron spectabile Steyerm.	Colombia, J. Rova et al. 2093 (S)	AY730297	AF242934	
Gonzalagunia affinis Standl. ex Steyerm.		AY730295		
		$AF323061^{2}$	AF242963	
Guettarda uruguensis Cham. & Schltdl.	cultivated at Fairchild Tropical Garden, FTG x 5-127, Gillis 9575 (FTG)	AY730294		
Hodykinsonia ovatiflora F. Wuell.	Australia, Puttock 8602667 (UNSW)	AY730293		
Luculia grandifolia Ghose		AJ3468963		
Mazaea phialanthoides (Griseb.) Krug & Urb.	Cuba, J. Rova et al. 2264 (GB)	AY730302	AF242980	AF1527496
Mazaea shaferi (Standl.) Delprete	ITS: Cuba, T. McDowell 4826-1993 (DUKE); rps16: Cuba, Rova et al. 2224 (GB)	AY730304	AF242911	AF152750 ⁶
Phyllomelia coronata Griseb.	Cuba, P. G. Delprete et al. 8913 (UPS)	AY730303		
Rachicallis americana (Jacq.) Hitchc.	cultivated at Fairchild Tropical Garden, FTG 64-266, Fanning KF81 (FTG)	AY730301	AF0040734	AF1527476
Rogiera amoena Planch.	ITS: Mexico, Mexia 8982 (GB); rps16: Guatemala, Hawkes et al. 1962 (S)	AY730286	AF243000	AF1024747
Rogiera cordata (Benth.) Planch.	ITS: Guatemala, Gustafsson & Fredriksson 126 (GB); rps16: cultivated at Royal Botanical Garden, Kew, Chase 2267 (K)	AY730285	AF242999	AF1527156
Rogiera cordata (Benth.) Planch.	Guatemala, J. J. Castillo & R. Luarca 2005 (NY)			
Kovaeanthus strigosus (Brandegee) Borhidi Rovaeanthus suffrutescens (Brandegee) Borhidi	cultivated at PIBG, D. Lorence 8920 (FIBG) cultivated at the Bergius Botanical Garden, Stockholm, Bremer 2712 (S)	AY730291 AY730290	AF243003	AF1527386
	Cuba, P. G. Delprete et al. 8725 (UPS)	AY730306		
Rondeletia alaternoides subsp. brachyloba M. Fernández Zeq. &	Cuba, J. Rova et al. 2228 (GB)	AY730310	AF243006	$AF152740^{6}$

Table 1. Continue

		Ge	GenBank accession	number
Taxon	Voucher data	ITS	2 squ	trnL- F
Rondeletia apiculata Urb.	Cuba, J. Rova et al. 2243 (GB)	AY730312	AF243007	
	Dominican Remiblic P C Delinoto et al 7534 (IIPS)	A V 730391		
		170000 TT		
Kondeletia berteroana DC.	Dominican Republic, F. G. Delprete et al. 1501 (UFS)	A Y (50522		
Rondeletia chamaebuxifolia Griseb.	Cuba, Machado et al. s.n. 1995-03-16 (S)	AY730327		
Rondeletia cincta Griseb.	Jamaica, P. G. Delprete et al. 7503 (UPS)	AY730323		
Rondeletia sp.	Jamaica, P. G. Delprete et al. 7479 (UPS)	AY730324		
		AY730305	AJ7867655	
Rondeletia hameliifolia Dwyer & M. V. Hayden	Panama, J. H. Kirkbride & S. M. V. Hayden 164 (NY)	AY730326		
	ITS: Cultivated at PTBG, D. Lorence 8796 (PTBG); rps16: Puerto Rico,	AY730315	AF243012	AF152745 ⁶
	Acevedo-Rodriguez et al. 7691 (NY)			
Rondeletia intermixta Britton subsp. intermixta	Cuba, J. Rova et al. 2245 (GB)	AY730311	AF0040774	AF1527426
Rondeletia lomensis Urb.	Cuba, J. Rova et al. 2216 (GB)	AY730313		
Rondeletia miraflorensis M. Fernández Zeq. & Borhidi	Cuba, J. Rova et al. 2217 (GB)	AY730314	AF243009	
Rondeletia nipensis Urb.	P	AY730330		
Rondeletia nipensis subsp. moaensis		AY730325		
Rondeletia ochracea Urb.	Dominican Republic, W. C. Holmes et al. 6618 (NY)	AY730316		
Rondeletia odorata Jacq.	cultivated at the Bergius Botanical Garden, Stockholm, Bremer &	AY730307	AF243010	AF1527416
	Andreasen 3504 (UPS)			
Rondeletia pachyphylla Krug & Urb. subsp. myrtilloides M. Fernández Zeq. & Borhidi	Cuba, J. Rova et al. 2232 (GB)	AY730331	AF243011	
letia pachyphylla K	Cuba, P. G. Delprete et al. 8674 (UPS)	AY730317		
Rondeletia pilosa Sw.	evedo-R	AY730332	AF243014	AF1527446
Rondeletia pitreana Urb. & Ekman	Liogier 13966 (NY)	AY730289		
Rondeletia plicatula Urb.	Cuba, P. G. Delprete et al. 8716 (UPS)	AY730318		
Rondeletia portoricensis Krug & Urb.	ico, C. M. Taylor 11687	AY730333	AF243015	AF152743
Rondeletia purdiei Hook. f.	Ecuador, Corbisier-Baland 1905 (UPS)	AY730328		
Rondeletia stipularis (L.) Druce	Jamaica, P. G. Delprete et al. 7472 (UPS)	AY730319		
Rondeletia subcanescens M. Fernández Zeq. & Borhidi	Cuba, P. G. Delprete et al. 8761 (UPS)	AY730320		
Rondeletia subcanescens M. Fernández Zeq. & Borhidi	9	AY730329		
Stevensia minutifolia Alain	Dominican Republic, P. G. Delprete et al. 7540 (UPS)	AY730308		
Stevensia minutifolia Alain	Dominican Republic, A. Liogier 13663 (NY)	AY730309		
Suberanthus brachycarpus (Griseb.) Borhidi & M. Fernández Zeq.	Cuba, T. McDowell 4824-1993 (DUKE)	AY730300		
Timonius nitidus FernVill.		$AF323063^{2}$		

The cited literature for published sequences includes: ¹McDowell et al., 2003; ²Moynihan & Watson, 2001; ³Razafimandimbison & Bremer, 2002; ¹Struwe et al., 1998. Sequences were also taken from ⁵S. Stranczinger, F. Jakab, J. L. Szentpeteri & A. Borhidi, unpublished.

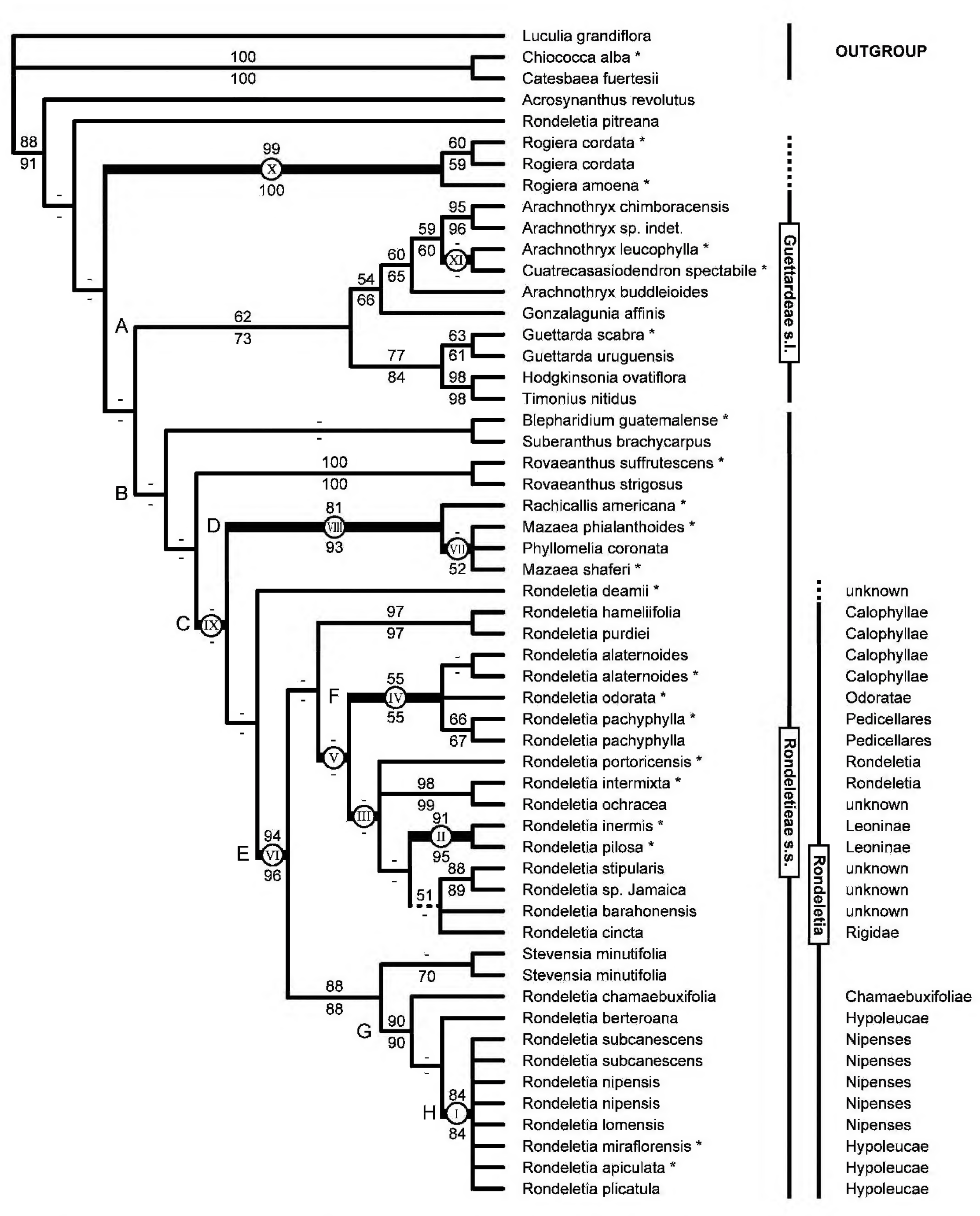


Figure 1. Tree compiled from the strict consensus trees from the two heuristic searches, without and with indels coded, respectively. The one dotted branch was not found in the heuristic searches, only in the jackknife search without indels coded. Numbers indicate jackknife support: numbers above branches are support values without indels coded, and numbers below branches are support values with indels coded. Jackknife support of 50 and lower is indicated by dashes (–). Letters A–H indicate the clades discussed in the text: —A. Clade comprising representatives of the tribe Guettardeae. —B. Clade corresponding to Rondeletieae sensu Rova et al. (2002). —C. Clade comparable to the one in which Acrosynanthus was found in Rova et al. (2002). —D. Clade comprising Mazaea, Phyllomelia, and Rachicallis. —E. The Rondeletia s. str. clade. —F. Clade including representatives of sections Odoratae, Pedicellares, and the paraphyletic section Calophyllae. —G. Clade distinguished by having 1- to 3-flowered inflorescences and retrorse-pilose flowers (corresponding to sections Chamaebuxifoliae, Hypoleucae, and Nipenses). —H. Clade comprising Cuban representatives of sections Hypoleucae and Nipenses. Taxa marked by an asterisk (*) are included in the combined ITS, rps16, and trnL-F analysis presented in Figure 2. The circled Roman numerals I–XI on bold branches refer to clades in the combined ITS, rps16, and trnL-F analysis (Fig. 2).

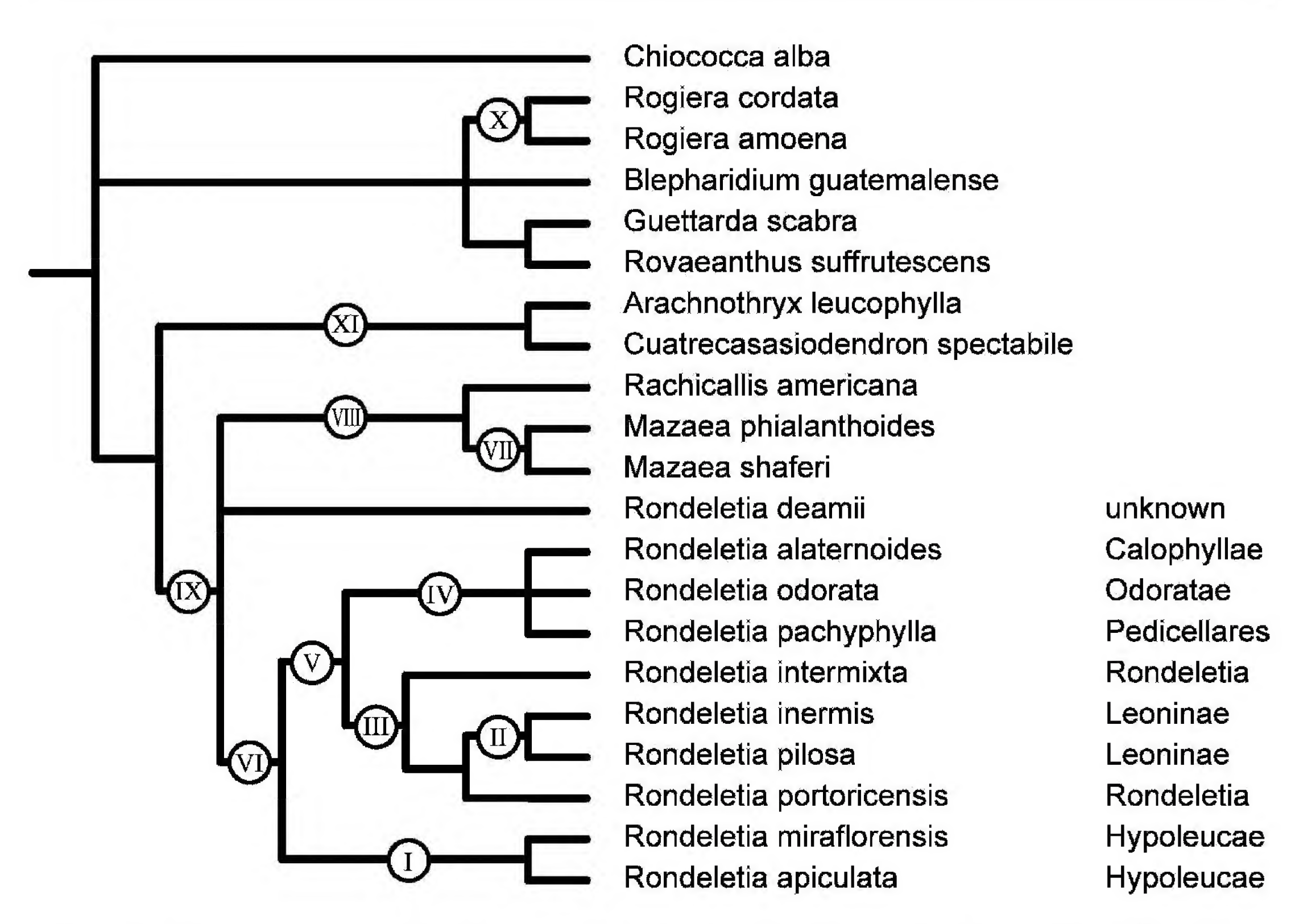


Figure 2. Strict consensus tree of the 12 equally parsimonious trees from the analysis of the combined ITS, rps16, and trnL-F data matrix. Guettarda scabra, Cuatrecasasiodendron spectabile, Rondeletia deamii, R. pachyphylla, R. miraflorensis, and R. apiculata were only represented by ITS and rps16 data in the data set. The circled Roman numerals I–XI on branches refer to clades in the tree in Figure 1. Sectional assignment of Rondeletia species, according to Fernández Zequeira (1994), is listed in the rightmost column. Rondeletia inermis and R. pilosa were not listed in Fernández Zequeira (1994), but have been assigned to section based on her key. Rondeletia deamii is not assigned to a section (cf. unknown among the sectional assignments).

Stevensia comprises 11 species endemic to Hispaniola. It is recognized by triangular stipules connected to a sheath, solitary and axillary flowers, two to three calyx lobes, five to seven stamens attached in the corolla throat, glabrous style, and ovoid to oblong seeds (Borhidi, 2001b). However, several of these character states are also found within *Rondeletia* s. str., according to Fernández Zequeira (1994). We therefore suggest that *Stevensia* should be included within *Rondeletia* s. str., pending future studies with an extended sampling.

GUETTARDEAE AND RONDELETIEAE

Our third aim was to compare a nuclear ITS phylogeny of the Rondeletieae with the results from a previous trnL-F chloroplast DNA (cpDNA) study

(Rova et al., 2002). The fourth aim was to see if ITS sequence data would place *Hodgkinsonia* in Guettar-deae or elsewhere.

Acrosynanthus revolutus Urb. and Rondeletia pitreana Urb. & Ekman (not classified to section) appear as early diversified lineages in the cladogram. One possible explanation for the position of R. pitreana could be that we were not able to read the sequence in its entirety. Because of this, it is about 40 bases shorter than the other Rondeletia sequences. Another possibility is that R. pitreana does not belong to Rondeletia. In any case, further studies are needed to solve the position of R. pitreana. The position of Acrosynanthus Urb. in the present analysis differs markedly from the results of the trnL-F study from Rova et al. (2002), where Acrosynanthus was found in a position equivalent to basal in clade C (Fig. 1). A

 \leftarrow

possible explanation would be that *Acrosynanthus* is not monophyletic: *A. latifolius* Standl. was included in the *trnL-F* study, but *A. revolutus* was sequenced in the ITS analysis. However, the possible paraphyly of *Acrosynanthus* must be left to another study when more material of this genus is available.

We found that the well-supported Guettardeae s.l. and Rondeletieae s. str. clades in the trnL-F study from Rova et al. (2002) have only weak support from ITS sequence data.

With regard to the tribe Guettardeae, there is moderate support for a clade including Arachnothryx, Gonzalagunia, Cuatrecasasiodendron, Guettarda, Hodgkinsonia, and Timonius (Fig. 1, clade A). According to our results, Cuatrecasasiodendron should be synonymized with Arachnothryx, and this is also morphologically supported (see taxonomic treatment below). While trnL-F data (Rova et al., 2002) showed Rogiera amoena Planch. and R. cordata (Benth.) Planch. as members of the Guettardeae, the inclusion of Rogiera s. str. in the Guettardeae clade is not supported by ITS data alone. In the combined analysis (Fig. 2), Rogiera is found within Guettardeae, while Arachnothryx is found to be more closely related to Rondeletieae.

In a recent molecular phylogenetic study, Achille et al. (2006) supported the monophyly of the Guettardeae as recognized here, although they showed that Guettarda, Antirhea Comm. ex Juss., and Stenostomum C. F. Gaertn. are polyphyletic. However, more genera and more species need to be included in the study in order to test the monophyly and delimitation of this tribe.

Although Rondeletieae sensu Rova et al. (2002) is recognized by ITS data in the consensus trees (Fig. 1, clade B), there is no jackknife support for this clade.

In both heuristic ITS searches, the genera *Blepharidium* Standl. and *Suberanthus* were found basal in the Rondeletieae s. str. clade, but again, there is no jackknife support for this. However, this position corresponds to the results from the *trnL-F* study of Rova et al. (2002).

The ITS data place *Rovaeanthus strigosus* (Benth.) Borhidi in the Rondeletieae, as sister taxon to *R. suffrutescens* (Brandegee) Borhidi. Just as in the *trnL-F* study (Rova et al., 2002), the ITS data indicate that *R. suffrutescens* belong to the Rondeletieae s. str. (although this is contradicted in the combined analysis where *R. suffrutescens* is found as sister to *Guettarda*). In any case, *R. suffrutescens* always appears in a separate position from *Rogiera*, and our study thus supports the transfer of these species from *Rogiera* into a new genus as proposed by Borhidi et al. (2004).

There is strong support for a close relationship between *Rachicallis DC., Mazaea* Krug & Urb., and

Phyllomelia Griseb. (D in Fig. 1). Rachicallis and Phyllomelia are monotypic genera, and Mazaea only comprises two species. Based on ITS data, one could argue that all three genera should be merged together. However, both Mazaea and Phyllomelia are easily distinguished by the peculiar fruit (pseudosamara, sensu Delprete, 1999b) and calyx morphology (Delprete, 1999b), and for this reason we prefer to regard them as separate genera.

HODGKINSONIA

The fourth aim was to see if ITS sequence data would place *Hodgkinsonia* in Guettardeae or elsewhere. Our study undoubtedly places *Hodgkinsonia* close to *Timonius*, which means within Guettardeae. This position is in accordance with the view of Mueller (1861) in the original description and Bremer (1992), but contradicts the supposition of Delprete (1996), who tentatively included the genus in tribe Chiococceae in agreement with Robbrecht (1988).

Conclusion

The ITS sequence data support only one of Fernández Zequeira's (1994) Rondeletia sections as monophyletic: section Leoninae. Rondeletia sections Calophyllae and Rondeletia are paraphyletic according to our analysis. However, one should bear in mind that we were not able to sequence more than one species from several sections. When we compare our ITS phylogeny with the character lists in Fernández Zequeira's treatment of *Rondeletia*, we were unable to find morphological characters that correspond with our phylogenies. The sections described by Fernández Zequeira are often defined by various combinations of overlapping character states, which makes comparisons difficult. The only exception is a clade including representatives from Rondeletia sections Chamaebuxifoliae, Hypoleucae, and Nipenses, which could be distinguished by having 1- to 3-flowered inflorescences and retrorse-pilose flowers. This clade could potentially be recognized as one section.

Rondeletia s. str. (i.e., excluding Arachnothryx, Javorkaea, Rogiera, Roigella, Rovaeanthus, and Suberanthus) has strong support, although some species need to be further investigated for their generic affinity (e.g., R. pitreana and R. deamii).

An important result from our study is that *Stevensia* minutifolia is included within *Rondeletia* s. str. A reevaluation of the morphological characters in *Rondeletia* (including *Stevensia*) based on the results from ITS and other sequence data is certainly needed. The present analysis clearly suggests that *Stevensia* should be recognized at most as a section of *Rondeletia*.

There is strong support for a division of the Rondeletieae-Guettardeae complex into the tribes Rondeletieae s. str. and Guettardeae s.l. from trnL-F data (Rova et al., 2002), but only weak support from ITS data. While part of the Guettardeae has moderate support, support for Rondeletieae in the sense of Rova et al. (2002) is weak in the ITS study, although the Rondeletieae s. str. clade is found in the strict consensus trees of all of our analyses, both including and excluding indel codings. It was not possible to compile a sufficiently large data set in order to test the delimitations of Guettardeae and Rondeletieae using a combined ITS, rps16, and trnL-F sequence data matrix; however, we consider that the phylogenies available up to this point (Rova, 1999; Rova et al., 2002; Delprete & Cortés-B., 2004; the present study) provide sufficient support for a re-delimitation of the tribe Rondeletieae.

Based on the results from ITS sequence data, we also reconsider Delprete's (1996) tentative inclusion (based on morphology) of *Hodgkinsonia* in the Chiococceae, since the present ITS sequence data support Bremer's (1992) conclusion (also based on morphological data) that *Hodgkinsonia* is part of the tribe Guettardeae.

TAXONOMIC TREATMENT

Based on the results from the present and other recent studies (Delprete, 1999b; Rova, 1999; Rova et al., 2002; Delprete & Cortés-B., 2004; Borhidi et al., 2004), we propose the following taxonomic descriptions and rearrangements.

Tribe Rondeletieae (DC.) Miq., Flora Nederl. Indië 2: 130, 156. 1856. Rondeletiinae DC., Prodr. 4: 342, 401. 1830, as subtribe "Rondeletieae," tribe Hedyotideae. Rondeletieae DC. ex Rchb., Der Deutsche Botaniker 1: 77. 1841, stat. non indic. TYPE: Rondeletia L.

Shrubs or trees; wood whitish or yellowish; raphides absent; axillary thorns absent. Stipules free or connate at base, mostly entire, rarely bifid, mostly interpetiolar, frequently with colleters on the adaxial side secreting resinous compounds, persistent to readily caducous; leaves opposite or verticillate, decussate, petiolate to sessile, blades chartaceous to thick-coriaceous; domatia variably present or absent. Inflorescences terminal or axillary, cymose, paniculate or thyrsoid, multiflorous or pauciflorous, or uniflorous. Flowers hermaphroditic, mostly actinomorphic, (3- to)4- to 6-merous; calyx persistent or caducous; lobes often minute, sometimes foliose; calycophylls commonly absent or pterophyllous (green to greenish white), with all calyx lobes expanding into

a rotate pterophyll after anthesis and present in all flowers in *Phyllomelia*; corolla hypocrateriform or narrowly infundibuliform, orifice with annular thickening, white, cream-white, red, green, or yellow, membranous to fleshy; aestivation valvate, contorted, or imbricate; stamens mostly as many as corolla lobes, inserted near the base or at the medial zone or near the orifice of corolla tube; anthers included or exserted, oblong to narrowly elliptic to button-shaped, 2-locular, opening by longitudinal slits, dorsifixed near the base or around the middle, introrse; pollen released as monads, colpate or colporate, exine reticulate or foveolate (not echinate); style branches present, with stigmatic surface smooth to verrucate; ovary inferior (half-inferior in *Rachicallis*), bilocular, with a few to many ovules (1 to 2 in Mazaea) per locule attached to a central placenta, or exceptionally one ovule per locule basally attached (Phyllomelia). Fruits woody capsules, loculicidal or septicidal, or septicidal and loculicidal contemporaneously (Blepharidium, Mazaea), commonly dehiscing basipetally, or exceptionally pseudosamaras, indehiscent (Phyllomelia); placenta central, rarely apically incomplete, or shortly stalked; seeds horizontal, imbricate, peltate, and vertical, minute, 3- to 5-angular or dorsoventrally convex, not winged, wing concentric or bipolar (Blepharidium, Mazaea), or basally inserted, ellipsoid-ovoid and fleshy (*Phyllomelia*).

Genera included: Acrosynanthus, Acunaeanthus Borhidi, Komlodi & Moncada, Blepharidium, Glionnetia Tirveng., Habroneuron Standl., Mazaea, Phyllomelia, Rachicallis, Rogiera, Roigella, Rondeletia, Rovaeanthus, Spathichlamys R. Parker, Stevensia, Suberanthus.

The description and delimitation of the Rondeletieae here proposed are based on the results of the present study in combination with those of Rova (1999) and Rova et al. (2002). The description is basically a reduction of that proposed by Delprete (1999a), based on his wide circumscription of the tribe to include the Condamineeae and the Sipaneeae, which was produced primarily for the floristic treatment and not based on a comprehensive phylogenetic analysis.

Rova (1999) and Rova et al. (2002) demonstrated that the Condamineeae (except the subtribe Portlandinae, which belongs to the Chiococceae s.l.) should be transferred to the subfamily Ixoroideae, in a complex also including the Calycophylleae and the Hippotideae (more studies are needed to re-delimit these groups; Kainulainen & Bremer, unpublished). Delprete and Cortés-B. (2004) and Rova et al. (2002) also demonstrated that the Sipaneeae belongs to the subfamily Ixoroideae and is a monophyletic group that was positioned in the same clade as the tribes

Henriquezieae and Posoquerieae in their phylogenetic analysis (Delprete et al., 2004).

The monotypic genus *Rachicallis*, endemic to the Caribbean Basin, is added (not included in the tribe by Delprete, 1999a) to the present delimitation of the Rondeletieae, which was placed close to this tribe in Bremer et al. (1995) and shown to belong to Rondeletieae by Rova et al. (2002) and in the present study.

As a result of this study, *Stevensia* is perhaps best treated as synonymous with *Rondeletia*, because in the phylogenies obtained it is positioned within the *Rondeletia*. However, as only one species of *Stevensia* (*S. minutifolia*) was included in the analysis, we refrain from proposing the necessary new combinations.

TAXA TRANSFERRED TO THE TRIBE GUETTARDEAE

Steyermark (1964) positioned Cuatrecasasiodendron in the Rondeletieae because of its foliaceous calyx lobes, capsular fruits, horizontal seeds, ovary with many ovules in each locule, and corolla with imbricate lobes. At the same time, he treated it as closely related to Rondeletia because of the corolla lobes being subzygomorphic, as the most interior lobe is more pubescent internally than the external ones, and glabrous or almost glabrous externally, while the others are pubescent externally. This genus was maintained in the Rondeletieae by Delprete (1999a) because of the same characters as used by Steyermark. However, in the phylogenies produced in the present study, Cuatrecasasiodendron was found within the Arachnothryx clade of the tribe Guettardeae, and the two taxa are treated here as synonymous.

In addition, a detailed analysis of the two species of Cuatrecasasiodendron described by Steyermark was undertaken. Steyermark (1964) distinguished C. spectabile Steyerm. from the type species because of its leaf blades hirsute below (vs. adpressed-pilose to arachnoid-pubescent below), shorter petioles, shorter stipules, corollas 17–20 mm long (vs. ca. 28 mm long, with longer pubescence), and longer and more secundiflorous inflorescence branches among other characters. A comparison of the type specimens with recent collections revealed that the characters used by Steyermark to separate the two taxa fall into a morphologic (and geographic) gradient.

The types of both taxa of Cuatrecasasiodendron were collected in the Valle del Cauca Department (Colombia); however, C. spectabile is from a low elevation of the coastal region, while C. colombianum Standl. & Steyerm. is from higher elevations of the Central Cordillera, and recent collections showed intermediate characteristics. Therefore, the two spe-

cies are treated here as synonymous to one another, and only one new combination in *Arachnothryx* is necessary.

- **Arachnothryx** Planch., Fl. Serres Jard. Eur. 5: 442. 1849. TYPE: *Arachnothryx leucophylla* (Kunth) Planch. (≡ *Rondeletia leucophylla* Kunth).
- Cuatrecasasiodendron Standl. & Steyerm., Acta Biol. Venez. 4: 29. 1964. Syn. nov. TYPE: Cuatrecasasiodendron colombianum Standl. & Steyerm.
- Arachnothryx spectabilis (Steyerm.) Rova, Delprete & B. Bremer, comb. nov. Basionym: Cuatrecasa-siodendron spectabile Steyerm., Acta Biol. Venez. 4: 33. 1964. TYPE: Colombia. Valle del Cauca Department: Costa del Pacifico, Río Cajambre, Barco, 5–80 m, 21–30 Apr. 1944 (fl.), J. Cuatrecasas 17165 (holotype, US!; isotype, VEN!).
- Cuatrecasasiodendron colombianum Standl. & Steyerm., Acta Biol. Venez. 4: 30. 1964. Syn. nov. TYPE: Colombia. Valle del Cauca Department: Cordillera Central, Vertiente Occidental, Hoya del Río Achicayá, Quebrada El Retiro, 300 m, 19 Dec. 1942 (fl.), J. Cuatrecasas 13694 (holotype, F!; isotype, US!).

Additional specimen examined. COLOMBIA. Depto. Valle del Cauca: Mun. Buenaventura, rd. Queremal—Anchicayá, Km 35, ca. 03°37′N, 76°53′W, ca. 300 m, 9 Apr. 1994 (fl.), J. H. E. Rova, L. Andersson, C. Gustafsson & C. Persson 2093 (GB).

Literature Cited

- Achille, F., T. J. Motley, P. P. Lowry II & J. Jérémie. 2006. Polyphyly in *Guettarda* L. (Rubiaceae, Guettardeae) based on nrDNA ITS sequence data. Ann. Missouri Bot. Gard. 93: 103–121.
- Andersson, L. & J. H. E. Rova. 1999. The *rps16* intron and the phylogeny of the Rubioideae (Rubiaceae). Pl. Syst. Evol. 214: 161–186.
- Borhidi, A. 1982. Studies in Rondeletieae (Rubiaceae). III. The genera *Rogiera* and *Arachnothryx*. Acta Bot. Acad. Sci. Hung. 28: 65–71.
- ———. 1993–1994 [1994]. Studies in Rondeletieae. XII. New combinations of Mexican and Central American taxa. Acta Bot. Hung. 38: 139–142.
- ———. 2001a. Additions and corrections to the "Nomen-clator of Mexican and Central American Rubiaceae" of D. H. Lorence. Acta Bot. Hung. 43: 37–78.
- ——. 2001b. Revisión taxonómica del género *Stevensia* Poit. Acta Bot. Hung. 43: 287–298.
- ——— & M. Fernández Zequeira. 1981a. Studies in Rondeletieae (Rubiaceae) I. A new genus: *Roigella*. Acta Bot. Acad. Sci. Hung. 27: 309–312.
- ——— & M. Járai-Komlódi. 1983. Studies in Rondeletieae (Rubiaceae). IV. A new genus: *Javorkaea*. Acta Bot. Hung. 29: 13–27.

- Bremer, B. 1992. Phylogeny of the Rubiaceae (Chiococceae) based on molecular and morphological data—Useful approaches for classification and comparative ecology. Ann. Missouri Bot. Gard. 79: 380–387.
- ——— & M. Thulin. 1998. Collapse of Isertieae, reestablishment of Mussaendeae, and a new genus of Sabiceeae (Rubiaceae): Phylogenetic relationships based on *rbcL* data. Pl. Syst. Evol. 211: 71–92.
- Delprete, P. G. 1996. Evaluation of the tribes Chiococceae, Condamineeae, and Catesbaeeae (Rubiaceae) based on morphological characters. Opera Bot. Belg. 7: 165–192.
- ———. 1999a. Rondeletieae (Rubiaceae)—Part I. Fl. Neotrop. Monogr. 77: 1–226.
- ——— & R. Cortés-B. 2004. A phylogenetic study of the tribe Sipaneeae (Rubiaceae, Ixoroideae), using *trnL-F* and ITS sequence data. Taxon 53: 347–356.
- Doyle, J. J. & J. L. Doyle. 1987. A rapid DNA isolation procedure for small quantities of fresh leaf tissue. Phytochem. Bull. 19: 11–15.
- Fernández Zequeira, M. 1993–1994 [1994]. Estudio taxonómico del género *Rondeletia* L. s.l. (Rubiaceae) en Cuba. Acta Bot. Hung. 38: 47–138.
- Lorence, D. H. 1991. New species and combinations in Mexican and Central American *Rondeletia* (Rubiaceae). Novon 1: 135–157.

- McDowell, T., M. Volovsek & P. Manos. 2003. Biogeography of *Exostema* (Rubiaceae) in the Caribbean region in light of molecular phylogenetic analyses. Syst. Bot. 28: 431–441.
- Moynihan, J. & L. E. Watson. 2001. Phylogeography, generic allies, and nomenclature of Caribbean endemic genus *Neolaugeria* (Rubiaceae) based on ITS sequences. Int. J. Pl. Sci. 162: 393–401.
- Mueller, F. 1861. Fragmenta Phytographiae Australieae 2. Joannis Ferres, Melbourne.
- Popp, M. & B. Oxelman. 2001. Inferring the history of the polyploid *Silene aegaea* (Caryophyllaceae) using plastid and homoeologous nuclear DNA sequences. Molec. Phylogenet. Evol. 20: 474–481.
- Razafimandimbison, S. & B. Bremer. 2002. Phylogeny and classification of Naucleeae s.l. (Rubiaceae) inferred from molecular (ITS, *rbcL*, and *trnT-F*) and morphological data. Amer. J. Bot. 89: 1027–1041.
- Robbrecht, E. 1988. Tropical woody Rubiaceae. Opera Bot. Belg. 1: 1–272.
- Rova, J. H. E. 1999. The Condamineeae–Rondeletieae–Sipaneeae Complex (Rubiaceae). Doctoral Dissertation, Botanical Institute, University of Göteborg, Göteborg, Sweden.
- Standley, P. C. 1918. Rondeletia. N. Amer. Fl. 32: 3–86.
- Steyermark, J. A. 1964. Novedades en las Rubiaceas Colombianas de Cuatrecasas. Acta Biol. Venez. 4: 1–117.
- Struwe, L., M. Thiv, J. W. Kadereit, A.-R. Pepper, T. J. Motley, P. J. White, J. H. E. Rova, K. Potgeiter & V. A. Albert. 1998. *Saccifolium* (Saccifoliaceae), an endemic of Sierra de la Neblina on the Brazilian-Venezuelan border, is related to a temperate-alpine lineage of Gentianaceae. Harvard Pap. Bot. 3: 199–214.
- Swofford, D. L. 2002. PAUP*: Phylogenetic Analysis Using Parsimony (*and Other Methods), Version 4. Sinauer Associates, Sunderland, Massachusetts.